 Syndrome of Inappropriate Secretion of Antidiuretic Hormone (SIADH) and Gerhardt Syndrome Associated with Shy-Drager Syndrome

Hirohito Sone, Yukichi Okuda, Chieko Bannai, Seiji Suzuki, Takashi Yamaoka, Yukari Asakura, Yasushi Kawakami, Masato Odawara, Teruhiko Matsushima, Koichi Kawai, Toshihiro Yoshizawa*, Hidehiro Mizusawa* and Kamejiro Yamashita

This is the first report on a case of syndrome of inappropriate secretion of antidiuretic hormone (SIADH) associated with Gerhardt syndrome (paralysis of bilateral vocal cords). A 67-year-old Japanese man suffering from progressive autonomic failure was diagnosed as having Shy-Drager syndrome (SDS) with hyponatremia due to SIADH and severe sleep apnea caused by a bilateral recurrent nerve palsy. Water load test showed alteration in diuresis which was corrected by phenytoin. Arginine vasopressin secretion was not suppressed by plasma osmolality below 280 mOsm/kg H2O. Impairment of the afferent pathways of baroreceptors, or impairment of the osmoreceptors could be speculated as the etiological factor of the SIADH observed in this case. (Internal Medicine 33: 773-778, 1994)

Key words: arginine vasopressin (AVP), autonomic dysfunction, multiple system atrophy (MSA), phenytoin (diphenylhydantoin), pulse oxymeter

Introduction

Gerhardt syndrome is sometimes observed among patients with Multiple System Atrophy (MSA) including Shy-Drager syndrome (SDS). It is characterized by paralysis of both vocal cords and frequently sudden death (1–6). There are some reports of SIADH in patients with diabetic autonomic neuropathy (7, 8). However, SIADH with autonomic dysfunction in cases with MSA or SDS has not been reported. It was demonstrated that patients with MSA including SDS have an impaired arginine vasopressin (AVP) response when they are stimulated by upright posture despite the presence of severe orthostatic hypotension (9–12). Here, we report a case of SIADH and Gerhardt syndrome associated with SDS in a middle-aged man.

Case Report

A 67-year-old Japanese man was suffering from various autonomic symptoms from the age of 56. These consisted of constipation, urinary incontinence, disturbance of perspiration, impotence and severe orthostatic hypotension. During admission to the Tsukuba University Hospital, clinical and laboratory results showed moderate hyponatremia [serum sodium (Na) level of 124 mEq/l], slight cerebellar ataxia and oily masked face. Based on various autonomic function tests (Table 1), he was diagnosed as having SDS. His marked snoring and sleep disturbance were investigated by polysomnography which exhibited severe sleep apnea for up to three minutes. Laryngoscopy revealed incomplete paralysis of bilateral vocal cords. At the age of 64, a pacemaker was implanted due to sick sinus syndrome. The electrocardiogram did not show any signs of pacing failure. Conventional radiography and computed tomography (CT) scan of the chest did not show any active intrathoracic lesions. CT of the brain including hypothalamus and pituitary did not exhibit any abnormalities except slight diffuse cerebral atrophy and a few small lacunar infarctions, both of which are age-related alterations. Laboratory data (Table 2) showed a persistent and relatively high urine osmolality (approx. 500 mOsm/kg H2O) despite a low plasma osmolality (268 mOsm/kg H2O). The human atrial natriuretic peptide (hANP) level was twice the normal level. While the basal serum growth hormone (GH) level was high, the GH response to either insulin (0.1U/kg)-induced hypoglycemia or growth hormone releasing hormone (GRH) (1μg/kg) was markedly low (Table
Table 1. Results of Autonomic Function and Other Neurological Tests

Inclination test (table tilting 60 degrees over the horizontal):
- Aug. 1992: Blood pressure 135/86→95/66 mmHg (11 min.)
  AVP 0.48→0.56 pg/ml
- Dec. 1993: Blood pressure 125/54→88/55 mmHg (10 min.)

Hand cooling test:
- Aug. 1992: Blood pressure 137/76→110/71 mmHg
- Dec. 1993: Blood pressure 129/86→128/83 mmHg

Perspiration test:
- Almost no sweating in 38°C, 30% humidity

Pupil response tests (eye dropping of epinephrine, cocaine, tylamine):
- Normal response

Shirmer, Gum test:
- Normal response

Uroflowmetry:
- Voiding volume (VV) 51 ml
  Residual volume (RV) 160 ml
  Maximal flow rate (MFR) 4.2 ml/sec.

Cystometry:
- First desire to void (FDV) 175 ml
  Maximal desire to void (MDV) 206 ml
  Maximal voiding pressure (MVP) 125 cmH2O

Nerve conducting velocity:
- Sensory nerve (median n., peroneal n.) 56-63 m/sec.
  Motor nerve (median n., sural n.) 40-59 m/sec.

Table 2. Laboratory Data on the First (Aug. 1992) and Second (Dec. 1993) Admissions

<table>
<thead>
<tr>
<th></th>
<th>Aug. 92,</th>
<th>Dec. 93,</th>
</tr>
</thead>
<tbody>
<tr>
<td>White blood cell count</td>
<td>7,600, 7,500/μl</td>
<td>7,600, 7,500/μl</td>
</tr>
<tr>
<td>Red blood cell count</td>
<td>371, 389×10^6/μl</td>
<td>371, 389×10^6/μl</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>12.8, 12.7 g/dl</td>
<td>12.8, 12.7 g/dl</td>
</tr>
<tr>
<td>Platelet</td>
<td>22.2, 36.6×10^9/μl</td>
<td>22.2, 36.6×10^9/μl</td>
</tr>
<tr>
<td>Total protein</td>
<td>6.5, 6.0 g/dl</td>
<td>6.5, 6.0 g/dl</td>
</tr>
<tr>
<td>Albumin</td>
<td>4.3, 3.4 g/dl</td>
<td>4.3, 3.4 g/dl</td>
</tr>
<tr>
<td>Na</td>
<td>124, 138 mEq/l</td>
<td>124, 138 mEq/l</td>
</tr>
<tr>
<td>K</td>
<td>5.5, 3.6 mEq/l</td>
<td>5.5, 3.6 mEq/l</td>
</tr>
<tr>
<td>Cl</td>
<td>89, 97 mEq/l</td>
<td>89, 97 mEq/l</td>
</tr>
<tr>
<td>Ca</td>
<td>9.0, 8.6 mg/dl</td>
<td>9.0, 8.6 mg/dl</td>
</tr>
<tr>
<td>Blood urea nitrogen</td>
<td>14.8, 10.0 mg/dl</td>
<td>14.8, 10.0 mg/dl</td>
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<tr>
<td>Creatinine</td>
<td>0.9, 0.6 mg/dl</td>
<td>0.9, 0.6 mg/dl</td>
</tr>
<tr>
<td>Uric acid</td>
<td>3.3, 3.6 mg/dl</td>
<td>3.3, 3.6 mg/dl</td>
</tr>
<tr>
<td>Total cholesterol</td>
<td>133, 157 mg/dl</td>
<td>133, 157 mg/dl</td>
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<tr>
<td>Triglycerides</td>
<td>51, 80 mg/dl</td>
<td>51, 80 mg/dl</td>
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<tr>
<td>Aspartate aminotransferase</td>
<td>31, 13 IU/l</td>
<td>31, 13 IU/l</td>
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<tr>
<td>Alanine aminotransferase</td>
<td>49, 11 IU/l</td>
<td>49, 11 IU/l</td>
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<tr>
<td>Lactate dehydrogenase</td>
<td>433, 401 IU/l</td>
<td>433, 401 IU/l</td>
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<tr>
<td>Alkaline phosphatase</td>
<td>74, 116 IU/l</td>
<td>74, 116 IU/l</td>
</tr>
<tr>
<td>γ-Glutamyl transpeptidase</td>
<td>58, 31 IU/l</td>
<td>58, 31 IU/l</td>
</tr>
<tr>
<td>Cholinesterase</td>
<td>145, 186 IU/l</td>
<td>145, 186 IU/l</td>
</tr>
<tr>
<td>Total bilirubin</td>
<td>0.8, 0.3 mg/dl</td>
<td>0.8, 0.3 mg/dl</td>
</tr>
<tr>
<td>Amylase</td>
<td>146, 66 IU/l</td>
<td>146, 66 IU/l</td>
</tr>
<tr>
<td>C-reactive protein</td>
<td>0.1, 0.4 mg/dl</td>
<td>0.1, 0.4 mg/dl</td>
</tr>
<tr>
<td>Fasting plasma glucose</td>
<td>96, 109 mg/dl</td>
<td>96, 109 mg/dl</td>
</tr>
<tr>
<td>Glycosylated hemoglobin</td>
<td>6.5, 6.6%</td>
<td>6.5, 6.6%</td>
</tr>
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</table>

Osmolality:
- Plasma (Aug. 92): 268, 277 mOsm/kgH2O
  (Dec. 93): 485, 576 mOsm/kgH2O

Adrenocorticotropic hormone:
- Plasma (Aug. 92): 41.7 pg/ml
  (Dec. 93): 32.9 pg/ml

Thyroid stimulating hormone:
- Plasma (Aug. 92): 0.75 μIU/l
  (Dec. 93): 0.39 μIU/l

Growth hormone:
- Plasma (Aug. 92): 0.0 mg/ml
  (Dec. 93): 0.0 mg/ml

Follicle stimulating hormone:
- Plasma (Aug. 92): 6.5, 9.7 μIU/l
  (Dec. 93): 6.5, 9.7 μIU/l

Prolactin:
- Plasma (Aug. 92): 9.5, 6.5 ng/ml
  (Dec. 93): 9.5, 6.5 ng/ml

Luteinizing hormone:
- Plasma (Aug. 92): 6.1, 5.9 μg/dl
  (Dec. 93): 6.1, 5.9 μg/dl

Free thyroxine:
- Plasma (Aug. 92): 1.35 μg/dl
  (Dec. 93): 1.35 μg/dl

Cortisol:
- Plasma (Aug. 92): 6.8, 5.9 μg/dl
  (Dec. 93): 6.8, 5.9 μg/dl

17-hydroxycorticosteroids:
- Plasma (Aug. 92): 4.4, 3.7 mg/day
  (Dec. 93): 4.4, 3.7 mg/day

17-ketosteroids:
- Plasma (Aug. 92): 3.7, 3.7 mg/day
  (Dec. 93): 3.7, 3.7 mg/day

Plasma renin activity:
- Plasma (Aug. 92): 0.6, 0.2 μg/ml
  (Dec. 93): 0.6, 0.2 μg/ml

Aldosterone:
- Plasma (Aug. 92): 9.9, 7.6 ng/dl
  (Dec. 93): 9.9, 7.6 ng/dl

Human atrial natriuretic peptide:
- Plasma (Aug. 92): 119–154 pg/ml
  (Dec. 93): 119–154 pg/ml

Catecholamine:
- Plasma (Aug. 92): 0.05 ng/ml
  (Dec. 93): 0.05 ng/ml

Epinephrine:
- Plasma (Aug. 92): 0.06 ng/ml
  (Dec. 93): 0.06 ng/ml

Norepinephrine:
- Plasma (Aug. 92): 0.20 ng/ml
  (Dec. 93): 0.20 ng/ml

Dopamine:
- Plasma (Aug. 92): 12.0 μg/day
  (Dec. 93): 12.0 μg/day

Norepinephrine:
- Plasma (Aug. 92): 30.1 μg/day
  (Dec. 93): 30.1 μg/day

Dopamine:
- Plasma (Aug. 92): 430.0 μg/day
  (Dec. 93): 430.0 μg/day

3). The GRH stimulation test was repeated once a day for five times, but the GH response did not improve. The response of plasma adrenocorticotropic hormone (ACTH) and serum cortisol to insulin-induced hypoglycemia was slightly low (Table 3). The basal serum cortisol and the daily excretion of 17-hydroxycorticosteroids (17-OHCS) were within the normal range and no evidence of adrenal failure was detected.

The result of the oral water load (20 ml/kg) test (Fig. 1) revealed a reduced diuresis (excretion of only 47 percent of load volume after 4 hours). The intravenous administration of phenytoin (500 mg), carried out simultaneously with oral water load test, markedly increased the diuresis (excretion of 117
Cortisol 10.3 9.1 13.9 13.2 10.2 (µg/ml)

ACTH 27.8 20.6 21.7 33.1 17.6 (pg/ml)

GH 3.3 1.6 2.5 3.3 2.8 (ng/ml)

a. the serum Na level remained normal under the same water oral administration of dopaminergic hypertensors. However, hypotension was also aggravated requiring intravenous and

tination of only 30 percent of load volume after 4 hours). Orthostatic

1) and revealed a slight aggravation of urine excretion (excre-

tion of only 30 percent of load volume after 4 hours). Plasma AVP levels

decreased only slightly during the water load. On the other hand, AVP response (from 0.48 to 0.56 pg/ml) to the head-up

inclination (table tilting of 60 degrees over the horizontal) was

very low despite the occurrence of severe orthostatic hypoten-

sion (from 135/86 to 95/66 mmHg) (Table 1). Water restriction

(750 ml/day) ameliorated the serum Na concentration to within

the normal range (135–140 mEq/l). However, oral administration

of phenytoin did not affect the serum Na levels.

A tracheotomy was indicated due to the risk of sudden death observed in Gerhardt syndrome; it was not performed however as the patient did not consent to it. Careful follow-up of his sleep apnea syndrome was carried out at home using a pulse oxymeter. The oxygen saturation was controlled during sleeping time; the patient was awoken when the oxygen saturation was lowered to 85% or less. Once discharged from hospital, he was observed in the outpatient clinic and he was doing well for almost a whole year with pulse oxymeter checking three or four times a night. However, as he could not sleep well due to the pulse oxymeter alarms, he chose to stop using it without any consultation with us.

A month after he stopped using the pulse oxymeter, he was apneic at midnight and he was referred to our hospital in August 1993. Cardiopulmonary resuscitation and tracheotomy were performed at this time. He received mechanical ventilation for about two weeks. During the convalescent period, there was no particular alteration in the laboratory data (Table 2), except in the serum Na level, compared to that of the former admission data. Oral water load test (20 ml/kg) was again carried out (Fig. 1) and revealed a slight aggravation of urine excretion (excretion of only 30 percent of load volume after 4 hours). Orthostatic hypotension was also aggravated requiring intravenous and oral administration of dopaminergic hypertensors. However, the serum Na level remained normal under the same water restriction.

Discussion

On the first admission, persistent hyponatremia and low plasma osmolality were seen, despite a higher urine osmolality as compared to that of the plasma. Psychogenic polydipsia and pseudohyponatremia was ruled out by the anamnesis. The etiology of the SIADH was investigated. Etiological factors like intrathoracic lesions, ectopic secretion by malignant tumor and drug-induced SIADH were not compatible with this case. Results of the phenytoin test suggest that inappropriate hypersecretion of AVP originated from the pituitary, rather than from an ectopic malignant lesion (13, 14).

The plasma hANP level was twice that of the normal concentration in this case. Plasma hANP levels of patients with SIADH increase to 6-fold the normal levels, though there are many individual variations (15). This is considered to occur in response to water retention secondarily induced by SIADH, because no significant correlation between plasma AVP and hANP levels is found in SIADH patients (16). Further, there is no report showing that hypersecretion of hANP is the primary cause of hyponatremia in patients with SIADH. In the current case, the elevation of hANP levels was considered to be caused by his VVI-type pacemaker which was reported to increase the hANP levels to twice that of the normal levels (17). Moreover, implanted pacemakers do not induce AVP secretion and do not cause SIADH. Thus, the cause of the SIADH in this case was not related to the hANP or the pacemaker.

To date, only a single case of SIADH associated with SDS (but without paralysis of vocal cords) has been reported (18). In their case, atrophy of the hypothalamus detected by magnetic resonance imaging (MRI) was considered as the etiological factor of the SIADH. These lesions were not detected by CT scan in the present case. MRI could not be performed because of his pacemaker.

The relationship between plasma osmolality and the AVP level in the present case (Fig. 2) can be explained by a hypersecretion of AVP in response to a plasma osmolality below 280 mOsm/kgH2O, though the AVP response to a higher plasma osmolality was almost normal. On the other hand, the AVP response to orthostatic postural stimulation is reported to be low in patients with MSA including SDS as was seen in this case also, though it was found to be high in patients suffering from other causes of orthostatic hypotension (9–12). So far, only one case of diabetes insipidus with SDS has been reported (19). Cluster breathing was associated with that case, suggesting damage of the pontomedullary respiratory center. It is suggested that the poor AVP response to orthostatic postural stimulation in SDS patients is primarily due to integrated lesions of the afferent baroreflex pathways (12).

In the present case, the real etiology of the SIADH is not clear. The afferent pathways of baroreceptors of the left atrium and the carotid sinus that reach the hypothalamus through the vagus and nucleus tractus solitarius are known to be inhibitory tracts (20). Hence, it can be speculated that both SIADH and bilateral recurrent nerve palsy could simultaneously occur by impairment of these tracts. Another possible explanation for the
low AVP response to head-up inclination and exaggerated secretion despite low plasma osmolality is a dysfunction of the osmoreceptors of low plasma osmolality of the hypothalamus that was not detected by the CT scan. Hypothalamic lesion in this case could explain the concurrent existence of hypersecretion of basal GH and the low GH response to the stimulation tests. These are sometimes observed among patients with SDS or Parkinsonism, which is considered to be caused by an abnormality of the tuberoinfundibular dopaminergic system of the hypothalamus (20–22).

AVP is a hormone that controls the peripheral circulation through its antidiuretic and pressor effects. However, recent
investigations (23, 24) revealed that AVP exerts dual and direct effects on the central nerve system acting as a neurotransmitter for inhibiting or accelerating the sympathetic nerve activity. The complex interaction between AVP and the autonomic system is presently under elucidation.

The etiology of Gerhardt syndrome associated with SDS or MSA is still obscure. Although there is one report stating that treatment with L-dopa produced a demonstrable improvement (2), this was not effective in the present case; tracheotomy is considered the only reliable therapeutic approach. On the first admission, we used a pulse oxymeter to prevent sudden death during periods of sleep apnea by making him wake up. It seems to be an effective method and it is worthy to attempt if the patients’ understanding and cooperation can be obtained.

In summary, this is the first report on a case of SIADH and Gerhardt syndrome associated with SDS. The findings suggest that SDS may be a causal factor of SIADH.

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